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#### (57) Abstract

The invention provides specific adsorbents and methods for separating an unsaturated hydrocarbon from a mixture containing such hydrocarbon. The adsorbents and methods are useful for separating dienes from mono-olefins. The adsorbents are selected from: a) an ion-exchanged zeolite selected from the group consisting of zeolite X, zeolite Y, and zeolite LSX, said zeolite having exchangeable cationic sites, and a plurality of said sites having silver cation or copper cation present; b) type A zeolite having exchangeable cationic sites, a plurality of said A zeolite sites having an alkali metal cation or an alkaline earth metal cation present.

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#### METHOD FOR SELECTIVE ADSORPTION OF DIENES FIELD OF THE INVENTION

The present invention relates to a process, and specifically selected adsorbents, for selective adsorption to separate a diene from a mixture, particularly one containing mono-olefin.

#### BACKGROUND OF THE INVENTION

In the preparation of commercial hydrocarbon product, it is often necessary to separate the desired product from other hydrocarbons having similar boiling points. For example, butene is synthesized commercially by processes which yield butadiene mixed with 1-butene. Conversely, butadiene product may also be contaminated with butene. Depending on the process used, the mixture may include other C3 and C4 hydrocarbon components including but not limited to 2-butene, other n-butenes, n-butane, isobutane, and isobutylene.

Purifying the mono-olefin, 1-butene, m is particularly troublesome due to closeness of its boiling point to that of 1,3-butadiene. In order to increase the purity of 1-butene, it is necessary to separate it from other hydrocarbons. Ordinarily, fractionation alone is incapable of completely separating 1,3-butadiene to achieve desired purity of the 1-butene mixtures. these Presently, butadiene is separated from paraffins primarily by distillation with selective solvents by absorption using solutions of absorbents. Extractive distillation is relatively energy-intensive, complex and not economical. Selective absorption with metal salt solution involves a significant number of stages with recycling of streams between stages. This method has the disadvantage of being energy-intensive and requiring handling and recirculating of of solvent streams which themselves contain contaminants or are subject to to be Current processes degradation. for olefin\paraffin

35 separation have not been sufficiently selective W. Walley . S

economically achieve the desired result for purifying mixtures of unsaturated hydrocarbons. Therefore, there remains the need for an improved method and improved adsorbents for use smin-methods to effectively economically purify unsaturated hydrocarbons. That some the edge of the expression of the that parefugiant elegated for solective a bir blon of a billion ್ 'ಹಿನ್ಸ್ ' . ಇತ್ತು ಅರ್ಚುಗಳ ಅರ್ಥವಾಗಿ ಕಟ್ಟಿಕ ಮಾಡುವುದು ಕಟ್ಟಿ ಕಟ್ಟಿಕ ಬರುವುದು ಬರುವುದು ಬರುವುದು ಬರುವುದು ಬರುವುದು ಬರುವ ្នាក់ ក្រសួងការពេលនៅ នៅក្នុងដោយស្រាយស្រាយស្រាយក្នុងសម្ព័ធ្យ ស្រុះ ពេលនេះ កម្មភាព សេដ្ transaction purvides methods and sucception useful but the commendations among the detail and the confidence of the confidence ್ರಾಗ್ ನರ್ಣ ಮ ರಾಹಿರ್ಜನೆಗಳಿಗೆ ವಿಗೆಕ್ಕೆ ಅರ್ಜನೆಕ್ಟ್ ಗುರ್ನಿಸಿದುಗೆ ಮುರ್ಗೆ ಅರ್ಜನೆಗಳಿಗೆ ಪ್ರಾಗಿಸಿಗಳು ಪ್ರಾಗಿಸಿಗಳು ಪ್ರಾಗಿಸಿಗಳು charities o cono-quefin guin as butene "Brasae Alaman edu Mac and the State of the ांकार उनमें तुरुक्तृ-Ciefin उनक्तिरांकी ने रिपेंट लंडेरका प्रेचाकेत come elegatificações com l'edanasunai filo ciunea e un un entreport ರವಾಸಗಾ ಸೀರಾಹಿಸಿದ ಕರ್ಡಿ ಇತ್ತಿ ಸಂಪರ್ಧಿಕ ಕಾರ್ಯ ಸಾರಾಷ್ಟ್ರಿ ಕೂಡುತ್ತಿರು. ಇಂತ್ರವಾಗು ಪ್ರಕಾರ range to the control of the consequence of the life in a formation and the consequence ំ ១ ១ ១០៩៤៦ ០០០ វិទី១ ខុខ១៩១២(វៀស្តី១៩ មុទ្ធប្រទេសចិន្ត្រី២ មានការប្រជាព្រះ ಕರ್ಮರ ೨ ೧೯೯೯-೬-೧೯ ೯- ದಿರಿ ಮುಂಗಳ ಗ್ರಾಕ್ಟಿಸಿಕರೆ ಈ ರಾಣಿ ಗೆರವರ್ಗ ಅವಹಾಗಿ ಮುಂದು will the sure of the sure to during guilled a end deriv ್ರಾಹ್ಮ ಪ್ರಕರ್ಣಕ್ಕೆ ಬಡೆತಾಗಿ ಅವರ ಕ್ಷಾಹ್ಮಿಸಿಕ್ಕಾಗಿ ಕಟ್ಟಿ ಆರಾಹ್ಮಿಕ್ಕಾಗಿ ಕಟ್ಟ चक्रवंदर्भ । अधिकेत्रे <sup>१९९५</sup> , १०० व्यक्तिक विद्यार 製造 デオールを封備 in the same stage is the Al Loidence ារធំបាន ខែ ខេត្ត នៅខេត្តទទួក។ ် အမြဲသေးသန်းကို သည်။ ကြိုင်းသည် သောက်သာဆုသျှမှာမြေသည်။ ကြိုင်းသည်။ မေးသည်။ သည် သည် သည် လည်းသွေး နည်း 1.5 အား မက္စာမည္းမရီး (ပရစ္စစ္ကည္းနဲ့ edo (၆၀) ကုန္ခဲ့နဲ့ ၂၈ ကုန္း ျခန္းျခန္းကို ್ರವ ಮುಸೀರಾಗಣೆಗೆ ಇದೇ ಅರ್ಭಾವರಿ ಅನ್ನೇ ಜೀರಾಕ್ಷವಗ್ರಾಧಾರರು ಪ್ರಧ್ಯಾಗ ಸಮ್ಮ ಪ್ರಾಗಂತ್ರದ ಪ್ರತಿವರಿಕೆ ಕನ್ನು ಸಂಪ್ರ చేసిపేగే వరాణాడులకానకి ఎక్కువా. ఈ అయ్యానినికి స్వేహి Reservation of Latitude and a a la fatti bila matar ್ರಾಟ್ ಮರ್ಗ್ ಕೃತಕರು. ಇತ್ತೂ ಕುಡೆಯ ಕ್ರಾಮೀ ಕ್ಲಡ  $p_{12} = p_{12} = p$ ut. 97 mgm to the late of the tile ရာရ မေန်းရ ခုန်းသို့ ကြည်းကြို့ ကြောင့် ကျည်ပြုပြု မရှိသော မြှေ့ရွှင့်ကျနေ့ အကျောက်ကြွေ ම 12 මානයෙනුව මුල්ලිල් මුල්ලි අද දුම්ලේ දුම් පුළුදුරිල දෙදැකුට 10 වන ඉදිනි සිටිම්ල්ල්ල්ල්ල්ල්ල්ල්ල්ල්ල්ල්ල්ල් និង សមានជនិងពេល ១៩៤០ ១៩៤០ ១៩៤៩៩៩៤៣ ៩ក្នុងរួម ១៩៧ ១៩៩៤១១០ រូបនាងមានន ತ್ತರ ಕ್ಷಾರ್ಥಿ ನರ್ಧಿಕಾದರು ಇಳಿ ಸರ್ವಿಶ್ವತನ್ನು ಆರಸ್ತಿಕ್ಕು ದಕ್ಕಾರೆದು ಇಡಿ ಸರ್ಗಾರ್ಯ ಸಂಗ್ರಿಸಿಕ ರಾಜಿಕೊಂಡಿ ಮೊತ್ತೆಗಳಿಂದ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕೊಂಡಿದೆ. ಅವರ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿಕೆ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿಸಿಕಾಗಿಗೆ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕಾಗಿ ಕ್ರಮಿಸಿಕ වැදියවෙන්න සෑ මෙම සිට මිට් වෙම් සිසෙන්ට සහ වා විසිදු සෑ Come විද්යාස්ත වඩා වා වඩා සිටු TA SE GO B GOOD LO ETÉ. LA MODEL MILO

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SUMMARY OF THE INVENTION

The invention provides new methods for separating unsaturated hydrocarbons from a mixture comprising the unsaturated hydrocarbons a The invention provides adsorbents specifically selected for accomplishing the separation. The absorbents and separation methods are particularly useful for selective adsorption of a diene from mixtures containing the diene; and are very effective for separating dienes from mono-olefins. In one aspect the invention provides methods and adsorbents to separate butadiene, hexadiene and/or octadiene from hydrocarbon mixtures; particularly where the hydrocarbon mixture contains a mono-olefin such as butene, hexane and/or octene.

Diene and mono-olefin compounds are often found together as a result of industrial processing. The separation of dienes from mono-olefin is difficult to achieve due to the closeness of their respective boiling points. This difficulty is illustrated by considering 1,3-butadiene, which has a boiling point of 4.49C and 1-butene which has a boiling point of 4.49C and 1-butene

The invention in one aspect, is particularly suited to cause such separation. Other important diene\mono-olefin separations include separation of hexadiene from hexene and separation of octadiene from octene. In the process of the invention, the diene is separated from a mixture comprising the diene by contacting the mixture with an adsorbent which preferentially adsorbs the diene. This produces a non-adsorbed component and a diene-rich adsorbed component.

In one aspect, the adsorbent comprises an ion-exchanged zeolite X, zeolite Y, and/or zeolite LSX. The selected zeolite has exchangeable cationic sites, with silver cation or copper cation present at some or all of the exchangeable cationic sites. Substantial cation exchange is preferred so that at least half of the cationic sites of the ion exchange zeolite contain a copper or

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silver cation. It is preferred that the ion exchange be . 2 \*\*\*\*\* particles of substantially or essentially complete so that the silver or copper ion exchange level of the exchangeable ion content is substantial enough to change the characteristic. In the case of the silver ion-exchanged zeolite, desirably a silver ion exchange level of the which is a exchangeable ion content is at least 70%, more desirably at least 80%, most desirably at least 85%, preferably at least 90% and more preferably at least 95% Most preferably, 10 substantially all (i.e. 99%), of the exchangeable ion sites Alternatively, the Xzeolite, Y-zeolite or zeolite LSX is a copper ion-exchanged and the ion exchange be as v : Like bush complete as described immediately above with respect to the Description silver cation is That is a at least half of the cationic and sites of the copper ion exchanged zeolite are occupied by copper copper. In the case of copper ion exchanged zeolite, the same level of high exchange content applies, according to the progressive levels stated above for silver. www.20 preferred that essentially complete ion exchange occur at 10. 100 - whereby substantially all (i.e., 99%) of the exchangeable \* if I among cationic sites are occupied by copper cations.

The copper and silver ion exchanged zeolites provide a junique advantage in that diene\mono-olefin 25 separation is facilitated by formation of π-complexation bonds. Therefore the silver ion exchanged zeolite, when complexation bonds for releasibly retaining the targeted unsaturated hydrocarbon desired to be selectively removed.

By formation of π-complexation bond, the silver or copper ion exchange zeolite is able to retain the targeted pressure. Thereafter the silver or copper ion-exchanged pressure. Thereafter the silver or copper ion-exchanged pressure are changed to exchange the adsorbed targeted hydrocarbon when all 35 its either or both of temperature and pressure are changed to see cause description (release).

estellan de la

ed an element of a targeted unsaturated hydrocarbon compound is achieved by contacting destine and a dimixture containing the targeted compound with an absorbent which comprises a type A zeolite. The type-A be zeolite is usable in its calcium form. The type A zeolite sis ab may also be used for selective adsorption where metal dations seffected from the group of alkaline metal cation term of alkaline earth metal cation are present. Therefore, whose server the Type-A zeolite may be represented by the nominal 10 general formula  $M^{41}_{x}Z^{42}_{y}A^{43}_{b}[(AlO_2)_{12}](S1O_2)_{12}]$  wherein  $M^{41}$  is an alkali cation, Z'asis an alkaline earth cation, and A' is be and we are a tri-valent cation, the value of x is 0 to 12, the value es e. commof y'is o to 6, and the value of b is at minimum zero and at maximum less than the sum of a plus y, provided that: x oin ada 15 wid 4 2y 43b is 12. For charge balance the maximum walue of of that bis less than 4. alkaline metal cation is selected from lithium, sodium, of grade core potassium, Frubidium and Caesium and mixtures thereof and the alkaline earth metal cation is selected from the group consisting of beryllium, magnesium, calcium, strontium, Sales the barium and mixtures thereof. The trivalent cation is with the preferably a metal compound capable of a trivalent valence astilities is Condition: Examples include aluminum and boron.

invention are used in a method for separating an unsaturated hydrocarbon from a mixture by accomplishing adsorption at a first selected pressure and temperature and then accomplishing release of desorption by changing at absorption is achieved at a pressure greater than the desorption (release) pressure. Preferential absorption pressure may be as high as about 35 atmospheres or more; and the desorption pressure of preferential adsorption is a thought atmosphere or less.

Desirably, the pressure of preferential adsorption is a range of about 1 atmosphere to about 35 atmospheres; and most desirably about 1 to 2 atmospheres. Desirably, the

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pressure of release is in a range of about 0.1 atmospheres to about 5 atmospheres; and most desirably in a range of about 0.1 atmospheres to about 0.5 atmospheres. The temperature of preferential adsorption is desirably selected to be in the range of about 0°C to about 150°C; more desirably up to about 120°C and most desirably in a range of about 25°C to about 80°C. The temperature of release is desirably in a range of about 70°C to about 25°C, and most desirably about 100°C to about 120°C.

The invention provides substantial advantages over conventional methods for separating dienes from a mixture due to the effective and economical process and adsorbents provided by the invention.

objects, features and advantages of the invention include an improved method for separating diene from a mono-olefin in a mixture, and particularly for separating C4 to C8 diene from a mixture which includes corresponding C4 to C8 mono-olefin. Another objective is to provide new adsorbents used in such new separation method.

These and other objects, features, and advantages will become apparent from the following description of the preferred embodiment, claims, and accompanying drawings.

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သူတုိ့ မုိင္းေတြကို အေျပာန္ျပီး ဖြစ္ပြားျမွဴး စုတာမွာေတြကုိ မုိင္းျပည္သည့္ တို႔ မိုက္သည့္ အေျပာက္သည့္။ ကို လုိင္းေတြကို အလည္း အမည္းလုိ႔ ဖြစ္ပြားျမွဴး စုတာမွာေတြကုိ မုိင္းျပည္နယ္ေတြကို တို႔ မိုက္သည္ လုံးစုံးျပည္သည့ 本作。(A) [1] [1]

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# BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows line representations of structure: (a) Solidate cage; or truncated octahedron; (b) type A zeolite "unit cell"; (c) "unit cell" of types X and 5 Y, or faujasite; (d); cation sites in type A (there are eight I, three IF, and twelve III sites per unit cell); (e) cation sites in types X and Y<sub>C</sub> (16 I, 32 I', 32 II, 32 II', 48 III, and 32 III's sites per unit cell)

Figure 2 Shows the pure-component 10 isotherms of T,3-CH6 mand strC4H8 on NaY and NaLSX type

zeolites at 70 c. seed see level seed 1,3-C,H, and 1-C,H, on Figure 3 shows uptake curves for 1,3-C,H, and 1-C,H, on

Nay-type zeolites. shows at because a someonent equilibrium isotherms for 1,3 cit; and 1-Cit. Agy zeolite at 70°C and

25°C.

25°C.

Figure 5 shows the equilibrium isotherms of C2H4 on NaY

Wen chivord on a and Agy type zeolites at 70°C. Figure 6 shows low pressure equilibrium isotherms for 260 1,3-C,H, and I-C,H, on 5A zeolite at 25°C.

20 1,3-C,H6 and 1-C,H8 on right 7 shows uptake curves for 1,3-C,H6 and 1-C,H8 on Sr exchanged 5A and 5A zeolites measured at 70°C. s billioners. Decin

Figure 8 shows desorption curve for temperature driven release of  $1,3-C_4H_6$  from 5A zeolite.

Figure 9 shows equilibrium adsorption isotherms for 25  $C_4H_6$  on AgY and NaY at 120°C.

Figure 10 shows low pressure equilibrium isotherms for  $1.3-C_4H_6$  and  $1-C_4H_8$  on 5A at 70°C.

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# DESCRIPTION OF THE PREFERRED EMBODIMENTS

The invention provides improved methods and adsorbents for separating a diene from a mixture which includes such diene. The invention is particularly useful for separating dienes having four to eight carbons from corresponding mono-olefins having four to eight carbons. Improved separation methods are needed due to the close boiling points of the "C4: toxoC8 dienes and corresponding C, to C, mono-olefins, Selected examples are as follows with numerical values being the boiling points in degrees centigrade 1,22 and 1,3-butadiene respectively 10.8 and -4.4; 1-butene, 2-2-butene cis and trans, respectively -6.3, 3.7 and 0.9; 1,3-, 1,4-, 1,5- and 2,4hexadiene respectively 73, 65, 59,5, and 80; 1-hexene, 2hexene cis and transmand. 3-hexene cis respectively 63.3, 68.8, 68, 66.4, and 67.1; and 1,7- and g Nidoe.**15**a 2,6- octadiene at 113-8 and 418-20; and 1-, 2-, 3-, and 4octene in a range of 121.3%125:6; all according to the 62nd

20 To one embodiments there is provided a process for separating diene from a feed mixture comprising the diene and at least one other unsaturated hydrocarbon compound, where the process comprises contacting the mixture with a diene-selective adsorbent which is an ionilensin 25 exchanged zeolite. Desirably, the ion-exchanged zeolite is The lower lies a Y zeolite, X zeolite, and/or low silica X zeolite, (LSX). Most desirably, the exchangeable cationic sites of the zeolite are ion-exchanged with silver cation or copper cation. It is most preferred that the zeolite be a silver zeolite (Ag-zeolite). The character of these zeolites will be described more particularly below. In the case of type LSX zeolite, it is preferred that it have a silicon to It is most 10 hoverhed at aluminum atomic ratio of less than 1.2.

dollar sations a preferred that the ratio of Si/Al be about 1.

It is desirable that more than half of the ion exchange sites of the zeolites contain the silver cation in the case of Ag-zeolite. It is preferred that in the case . . . . . 7

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of Ag zeolite, substantially all of the cation sites are occupied by silver in a +1 oxidation state (+1 valence state) . In the case of Cu-zeolite, it is also desirable that more than half of the ion exchange sites contain copper ions. It is also preferred that substantially all of the sites are ion exchanged with copper in the case of copper exchanged zeolites where the copper is in the +1 valence or oxidation state.

rence or oxidation state. O Spring Insanother embodiment, the invention provides a process for causing the separation of diene from one or more other unsaturated hydrocarbons, particularly monoolefin as described above, using a type A zeolite. In this process the diene is separated from a feed mixture utilizing the type A zeolite having an alkali-metal cation or an alkalines earth metal cation at exchangeable cationic sites to cause the

adsorption of the diene.

Desirably: commercially available 5A zeolite is The 5A zeolite contains calcium (Ca) as the major used. Bascoid a Park cation. This means that for all the available cation sites of A-zeolites Ca is found at over half such sites. minor Cation also sodium. Therefore, the 5A zeolite has substantially and essentially all of its cationic sites occupied by Ca and Na, but primarily Ca. This 5A zeolite is also referred to herein as CaA zeolite.

In one embodiment, the type A zeolite optionally contains a mixture of cations besides the alkali metal and alkaline earth metal. Therefore, at some of the cation exchange sites of the zeolite, cations other than +1 and +2 will be present. The nominal formula of this zeolite is M<sup>+1</sup><sub>x</sub>Z<sup>+2</sup>yA<sup>-3</sup><sub>b</sub>[(AlO<sub>2</sub>)<sub>12</sub> (SiO<sub>2</sub>)<sub>12</sub>] wherein the value of x is 0 to 12 and the value of y, is 0 to 6, and the value of b is at minimum zero and at maximum less than the sum of x plus y, provided that: x x 2y +3b is 12. This is believed to provide a mixture of cations in this type A zeolite which is balanced. It is preferred that when the A zeolite contains tri-valent (+3) cations, the atomic amount of such

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tri-valent is less than the combined amount of the alkali and alkaline metal cations. It is most preferred that in the aforesaid nominal general formula, the alkaline earth metal is primarily Ca, providing Ca-A (5A) zeolite.

The method and adsorbents (sorbents) of invention are particularly suited for temperature swing adsorption (TSA) and pressure swing adsorption (PSA). process is effective to separate diene from a mixture containing one or more sother unsaturated hydrocarbons and particularly emono-olefin compounds. Before describing the invention and the use of zeolites of the invention sin TSA and PSA processes, more particular description of the physical characteristics of the zeolites of the invention is provided along with preferred ion exchange method. ಿಪ್ರಕರ್ಮ ಚೇವರ ಚರ್ವಿನಗಳ

constant Zeolites have historically been used adsorbents due their selectivity. Zeolites selectively adsorb molecules on the basis of the size and the shape of the adsorbate molecule and are therefore called molecular sieves. Zeolites have been known for use in selective hydrocarbons. adsorption of carbon monoxide and Crystalline zeolite Yww zeolite A and zeolite X described in USPN:3,130,007; 2,882,243; 2,882,244; eachsof: which is incorporated by reference in its entirety. Type 5A zeolite, and type 13X zeolite are described for nitrogen adsorption in USPN 5,551,257. Types 4A (Na-A) and 5A (Ca-A) zeolites, are described in USPN 3,785,122: Low silica X zeolite (LSX) having Si/Al ratio less than or equal to 1.25, desirably less than or equal to 1.2, and preferably about 1 is described in USPN 5,268,023. Each of the aforementioned patents is incorporated herein by reference in its entirety. Consistent with the features described in these patents, zeclite characteristics are below excerpted from "Gas Separation Processes" by R.T. Yang (1987; Butterworth Publishers). 7 0

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Zeolites are crystalline aluminosilicates of alkali or alkali earth elements such as sodium, potassium, and calcium, represented by the stoichiometry:

M<sub>x/n</sub> (AlO<sub>2</sub>) (SiO<sub>2</sub>) | zH<sub>2</sub>O (

where x and y are integers with y/x equal to or greater than 1, n is the valence of cation M, and z is the number of water molecules in each unit cell. are shown in Figure 1(b) and (c). The cations are necessary to balance the electrical charge of the aluminum atoms, each having a net charge of 1. The water molecules can be removed with ease upon heat and evacuation, leaving an almost unaltered aluminosilicate skeleton with a void fraction between 0.2 and 0.5. The skeleton has a regular structure of cages, which are usually interconnected by six rain more water sa windows in each cage. The cages can imbibe or occlude large amounts of guest molecules; in place of water. size of the window apertures, which can be controlled by fixing the type and number of cations, ranges from 3 Å to 10 A. The sorption may occur with great selectivity because of the size of the aperture (and to a lesser extent because of the surface property in the cages) - hence the name molecular sieve.

At least forty species of naturally occurring zeolites have been stounds to The principal chabazite, (Ca, Na<sub>2</sub>) Al<sub>2</sub>Si<sub>4</sub>O<sub>12</sub>(6, H<sub>2</sub>O); gmelinite; (Na<sub>2</sub>, Ca)  $Al_2Si_4O_{12}$  (6  $H_2O)$ ; mordenite; (Ca,  $K_2$ , Na<sub>2</sub>),  $Al_2Si_{10}O_{24}$  (6.66  $H_2O)$ ; levynite;  $CaAll_2Si_3O_{10}$  (5  $H_2O$ ); and faujasite, (Na<sub>2</sub>, Ca, Mg,  $K_2$ )OAl<sub>2</sub>Si<sub>4.5</sub>O<sub>12</sub> (7.H<sub>2</sub>O)... More than 150 types of zeolites have been synthesized; they are designated by a letter or group of letters - Type A, Type X, Type Y, Type ZSM, and so The commercial production of synthetic zeolites started with the successful development of low-temperature (25-100°C) synthesis methods using very reactive materials such as freshly coprecipitated gels or amorphous solids.

The primary structural units of zeolites are the tetrahedra of silicon and aluminum, SiO4 and AlO4. units are assembled into secondary polyhedral building

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units such as cubes, hexagonal prisms, octahedra, truncated octahedratos The silicon and aluminum atoms, located atothe corners of the polyhedra, are joined by a shared oxygen . The final zeolite structure consists of assemblages of the secondary units in a regular threedimensional crystalline framework. The ratio Si/Al is commonly one to five The aluminum atom can be removed and replaced by silicon in some zeolites, thereby reducing the number of cations; and the cations can also be exchanged. 10 7 The inner atoms in the windows are oxygen. The size of the windows depends, then, or the number of oxygen atoms in the . Will all all aring and four, Alivey Isix, reight, then, por twelve. aperture size, as well as the absorptive properties, can be same to refurther modified by the number and type of exchanged eations A description of the structures will be given only for the zeolites important in gas separation, Type A based and Type X sand Yo. (See Table, The tag to substitute

Type A. 2 The structural unit in Type A. 2 The structural unit in Type A. zeolite, as well as in Types X manda Y, is the truncated octahedron, shown in Figure: 1 (a): OThis unit is also called sodalite cage, as sodalite is formed by directly fusing the fourbuy 300 member rings of the units. Two The four-member rings of the socalite units cantalso be linked through four-member prisms, as shown in Figure 1(b), which is type A zeolite. 25 The unit cell of Type A zeolite, as shown in this figure, contains 24 tecrahydra, 12 AlO, and 12 SiO, When fully the central hydrated, 27 water molecules are contained in the central cage or cavity of the unit cell, and in the eight smaller sodalite cages. The free diameter in the central cavity is 30 ... 11.4 A, which is entered through six eight-member oxygenring apertures with a minimum diameter of 4.4 Å. There are twelve negative charges to be balanced by cations in each unit cell. The most probable locations for the cations are i O.E has dindicated in Figure 1 (d) w Trype I ds at the center of the The six member ring, thus at one of the eight corners of the cavity. Type II is at the eight-member aperture, directly obstructing the entrance. Type III is near the four-member ring inside the cavity. Type A zeolites are synthesized in the sodium form, with 12 sodium cations occupying all eight sites in F and three sites in II, plus one site in III. This is the commercial Type 4A zeolite, with an effective aperture size of 3.8 Å. The sodium form can be replaced by various other cations or by ashydrogen ion. The commercial Type 3A zeolite is formed by exchanging Na with K, resulting in a smaller effective aperture size due to the larger K. The aperture size of the sodium form can also be increased by exchanging Na with Ca or Mg<sup>12</sup>, since 2 Na are replaced by one bivalent cation. The form of the exchanged Ca<sup>12</sup> or Mg<sup>12</sup> is Type 5A with rather unobstructed and larger apertures.

Types X and Y: The skeletal structure of Types X and Y zeolites are the same as that of the naturally occurring faujasite . The sodalite units are linked through six-member prisms as shown in the unit cell in Figure 1(c) SY Each unit cell contains 1925(Si, Al)O4 tetrahedra. ne the number of the aluminum long per unit cell varies from 96 to 77 for Type X zeolite, and from 76 to 48 for Type Y zeolite grains framework has the largest central cavity end of volume of any known amounting to about 50% void fraction in the dehydrated form, the Lunit cell, when fully \*\* The third ated, acoutains approximately: 235 water molecules, 25 25 mostly in the center cavity. The aperture is formed by the ville is if twelve-member oxygen rings traith gas free clameter of approximately 7:4 A: Three major locations for the cations The locations are: center of cavity (IP and II'); and at the twelve-member aperture (III and III') ... The commercial 10% zeolite contains Ca+2 as the major cátion, and Natris the major cation for 13X zeolite. The distribution of Nat,  $K^*$  , Car, other cations, and  $H_2O$  in  $35^{-1}$  X and Y zeolites. The BET surface area, measured, with N<sub>2</sub> for zeolites falls in the range between 500 and 800 m<sup>2</sup>/g.

ေဆာမ်ကြီးသည်။ ခုနှစ်ချင်သည် မို့နှစ်သည်။ ကိုသည် ကောင်းကျော်သည် ကောင်းကျော်သည် ကောင်းကျော်သည် ကောင်းကျော်သည် ကေ

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In one aspect, adsorbents of the invention, as described below, achieve selective adsorption by means of the \$\pi\$-complexation mechanism. Separation by \$\pi\$-complexation is a subgroup of chemical complexation where the mixture is contacted with a second phase, which contains a complexing agent. The advantage of chemical complexation is that the bonds formed are stronger than those by van der Waals forces alone, so it is possible to achieve high selectivity and high capacity for the component to be bound. At the same time, the bonds are still weak enough to be broken by using simple engineering operations such as raising the temperature or decreasing the pressure.

The m-complexation generally pertains to the main group (or d-block) transition metals, that is, from Sc to Local Cu, Y to Ag, and La to Au in the periodic table. These metals or their ions can form the normal so bond to carbon and, in addition, the unique characteristics of the doubitals in these metals or ions can form bonds with the unsaturated hydrocarbons (olefins) in a monclassic manner.

This type of bonding is broadly referred to as m-complexation, and has been considered for gaseous hydrocarbon separation and purification using cumbersome liquid solutions.

25 conventional a complexation sorbents did not provide a large working capacity for the target olefin. The examples below show that the sorbents of the invention have superior selectivity for 1,3-butadiene adscrption. The examples show the new adsorbents (sorbents) usable with preferred PSA and TSA processes for very effective diene separation. The PSA process is effectively usable with the new sorbents of the invention. Here, multiplicity of periodic steady states is an important aspect of cyclic adsorption process. For pressure swing adsorption (PSA), multiple steady states exist for a fixed set of operating

multiple steady states exist for a fixed set of operating conditions, over a particular range of one or more of these operating variables (i.e., bifurcation variables). The

final stable state depends only on the initial condition (i.e., the perturbation variables).

There are a variety of commercial applications in which component separation is conducted by pressure swing adsorption (PSA) and temperature swing adsorption (TSA).

Other less commonly used processes are volume swing adsorption and concentration swing adsorption. The most commonly described are PSA and TSA systems.

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adsofbable component is passed through an adsorbent bed capable of selectively adsorbing the more readily adsorbed component at an upper adsorption pressure. The eless readily adsorbable component passes through the bed and is readily adsorbable component passes through the bed and is readily adsorbable to a lower desorption pressure for desorption of the more readily adsorbable component, and see that the readily adsorbable component, and see the readily adsorbable component, and see the readily adsorbable component, and see the readily adsorbable component.

\* The temperature swing adsorption processing the Euros and in adsorbent is exposed to the feed mixture for a period of ameans little time during which the more readily adsorbed component is Thereafter the flow of the feed is shut off from adsorbed. and the adsorbent bed and the adsorbent is exposed to a purge #25 Fluid, typically a gas, which strips the adsorbed component resimilars of from the bedrand regenerates the bed for further use. TOWN STRANGER adsorbed component is driven off from the bed by as lightless sheating withe adsorbent in the Vregeneration state. Therefore, the temperature of desorption is higher than the temperature of the adsorption portion of the cycle. 6 3 00 00 Details of PSA and TSA adsorption will not be repeated The states with the following United States price in Patents, each of which is incorporated by reference herein, 11177 Deliging it si wentirety: 9 5,268,023; 5,551,257; 5,554,208; ; 7 8**35**; 7 75,672,196,7 5,672,196,25 800,20 800,20 900,608 810,100

Ion exchange of zeolite is easily accomplished by mixing the zeolite in an aqueous solution of the metal

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spine. Do salt. The metablof the salt is the metal to be exchanged into the cationic site. The concentration of the solution is varied according to the desired level of ion exchange. Fig. 1. The ion exchanged recline is then removed by filtration from the aqueous solution and washed free of the soluble salts this batch ion exchange is widely used due to its simplicity: Essentially complete ion exchange can be it achieved by repeating the same procedure several times. By such repeated treatments over 20% of the exchangeable 10 cationic sites are consexchanged with Ag. Preferably, Sover 95% of such sitescare ion exchanged. The same high TO CONTROL OF Exchange his tobtainable with the copper ion at The Agreeolites of the invention were prepared by ion exchanging with assolution of AgNO, containing five 15 times the cationic exchange capacity of the zeolites. was done to assure complete ton exchange. The Cu-zeolites and the sinvention were prepared by ion exchanging with a solution of CuCl2 or Cu(NO3)2, followed by reduction of Cu+2 o or sit white Cutto Cutto and wolf fore of the early of soldier was

20 20 The The fellowing examples show new sorbents for the challenging separation of removing trace amounts of dienes (50 PPM) from a normal alphasolefin (NAO) stream. sorbents of the wexamples below demonstrate operability for a pressure swing adsorption (PSA) or temperature swing adsorption (TSA) mode. i feri e hils sur bi so

gar the the sorbents of the example were pre-selected and weak chemical complexation. The first type of interaction derives from the molecular sieving effects of certain substrates such as zeolites or carbon molecular sieves (CMS). However, zeolites were mainly considered for this application due to atheir high Henry's adsorption. For this approach; the small differences in the kinetic diameters of mono-olefin are exploited. specifications of various sizes into To notify To the pore-aperture of type A zeolites.

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As A 13 A 210 of the Components of the components TWIN involved in the examples, weak chemical and complexationbased sorbents that have demonstrated the capabilities of 1 1 4 2 577 ( 17) adsorbing olefin over paraffins are not necessarily suited 1 5 % for diene separation from Imono to leftin . In the separation of diene from mono-olefin, both adsorbates are olefinic in The state of the s the double bonds am Double bonds that alternate with single alidient discribonds are said to benconjugated; double bonds that are STAGE 10 222 separated by more than one single bond are said to be Fisolated. A third, eless important class of dienes contain cumulated double bonds, that is, double bonds adjacent one were fanother and known assaltenes of the most common dienes is 1,3-butadiene ... Its boiling point is -4.4°C and This 15 dery close to 1 soutened 26 3°C) Separation of these two The imajor difference between the two is the This is the conjugate wdiene natures of 1,3-butadiene. Since 1,3 Date to butadiene is present in such low concentrations, the bond sensible to adsorbesignificant amounts of butadiene at such - That islow partial pressures lament is month extra one

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of the sorbent can be tailored by selecting specific cations and substrates. When dealing with bulk separations, high amounts adsorbed at low pressures are usually not desired, and are usually avoided because they limit the working capacity of the adsorbent. However, for this particular application, a high Henry's law constant is particular application, a high Henry's law constant is required. Therefore, zeolite based sorbents are a good candidate for substrates. It is shown that Ag' exchange Y zeolite has high affinity for olefins at very low pressures. Therefore, the examples focus on using zeolitic materials such as Type A. X. or Y. ion exchange with Cu' or Ag cations to carry out the separation of 1, 3-butadiene from 1-butene. It should be noted that the separation of the conjugated butadeine from butene is merely exemplary.

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The processes described below are also applicable to separate other dienes such as hexadiene and octadiene from mixtures containing hexene and octene. The temperatures and pressure given below are also exemplary.

Depending on the chosen conditions, the separation is accomplished in the gas phase, the liquid phase or some combination thereof, including for example, saturated gas. It should be noted that the butadiene and butene of the examples share common chemical characteristics with other diene/mono-olefin combinations described earlier. Therefore, the effectiveness of the invention in separating all such stated combinations is apparent.

The sorbents described in the examples below contained various cations included in high surface area substrates. The dispersion of cations was accomplished using ion exchange. The chemistry of ion exchange in zeolites is well documented. See USPN 2,882,243, 2,882,244 and 5,268,023, each of which is incorporated 20 herein in its entirety. All exchanges were performed similarly. They involved vacuum filtering and washing of the zeolite with deionized water. Compared to the original cation exchange capacity (CEC), each solution contained 10fold cation equivalents. This procedure ensured 100% exchange. For A (Linde), LSX (low silica X, UOP), and Y (Strem Chemicals) zeolites, the starting forms contain Na\*. The zeolites used were powder form (binderless). Prior to use the samples were calcined in vacuo at 350°C.

In the examples below, the hydrocarbons used as the adsorbates were 1,3 butadiene (CP grade, Matheson), 1butene (CP grade) and helium (pre-purified grade, Metro The gasses were used without further purification. Isotherms and uptake rates were measured 35 utilizing both a Shimadzu TGA-50 microbalance system following well known procedures. For low pressure data (<0.05 atm), a Micrometrics ASAP 2010 pore size

distribution analyzer was utilized. This system is capable of accurately measuring pressures in a range between 1 x 10-7 - 1.25 atm. While one can obtain data at very low pressures with the Micromeritics ASAP 2010, this is very time consuming. The Shimadzu TGA-50 thermogravimetric analyzer is used to screen potential sorbents. Also, measurements were made at two temperatures (25°C, 70°C) in order to obtain isosteric heats of adsorption. The overall diffusion time constants, D/R<sup>2</sup>, were calculated from the 10 uptake curves measured from a stepped pressure increment from 0. to 0.6 atm by well known methods and assumptions.

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Example 1

Pre-screening of Y-type Zeolites for 1,3-Butadiene Selectivity

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15 The framework structure of LSX and Y-type zeolites resembled that of the naturally occurring faujasite (see Figure 1). The pores of these zeolites are made of 12-member oxygen rings. They have an aperture size of about 8.1 A. They only differ in the ratios of Si/Al. 20 This ratio determines the number of cations per unit cell and therefore has a significant effect on the adsorption properties. Y-type zeolites contain between 56 and 76 cation sites, respectively. It is because of the properties mentioned above that these zeolites were selected as substrates. The large size facilitates the diffusion of large molecules such as 1,3butadiene and 1-butene. Also, the large number of cation sites available for exchange with Cu1+ or Ag1+ helps increase the adsorption capacity.

The pure-component equilibrium isotherms of 1,3-C4H6 and 1-C4H8 on NaY and NaLSX type zeolites at 70°C are shown in Figure 2. The partial pressures ranged from .06 to 1 atm. While these pressures are outside the range of interest for 1,3-C4H6, they are useful for screening potential substrates and for determining capacities for 1-

C4H3. Themequilibrium data was fitted with the Langmuir isotherm model: Thesequilibrium amounts of 193-C.H. and 1to the transfer of CaHa adsorbed at \$70° Chatqleatm on NaY zeolite were 2.80 and 2.69 mmol/g respectively. This equals about 36 molecules 5: 27 5 do of 1,3-C4H6 and 34, molecules of 1-C4H8 per unit cell of NaYmake curves for an atype zeolite. EUptake curves for a, 3-C, H, and 1-C, H, on Naytype zeolites are shown in Figure 3. They were obtained by change in pressure from 0-0.06 atm. Both components diffused very bfast to Complete uptake was # 10 streachieved within 50 seconds # This was due to the very large Topore aperture usize hoff the NaY zeolite, about 7.4 A. Diffusion time constants, D/R2, were calculated by fitting talest and experimental adata swith with solution of the diffusion as a sussit , equation for spherical particles as described by Karger and Ruthwen in 1992: The values of D/R2 obtained for 1,3-C4H6 ennagency wo and 1sC, He at 270°C, Were 70850 x 403 and 5.76 x 103 1/s, heardingue respectively reather ratios of sthe diffusivities is rather This is a secondary (and 3) for kinetic separation application. However, . We incorred favorable scharacteristic sof W-type zeolite is its 20 . 20 relatively high Henry's Law constant Just his indicates that the Yetype szeolites have good affinity for C4's and are The continue also good candidates for Constand Carstillings ាស់ 🔍 👉 សហជាមិស្ ៩២០ រៀបចំ ក្រសួងស ៩ ឬក្សាមេ ការ បកក្នុង បាស់ជាមេរិយកដា។

#### Example 2

As per Example 1, Y-type zeolites have a very product of high affinity for C4? stat low pressures. Therefore, this material is a good substrate for T-buttene purification, and also for shexene and octene purification. In order to increase the affinity of the NaY zeolite for 1,3 butadiene, the NaY cation was fully exchanged by Agt which is capable of m-complexation with olefinic compounds. The pure component equilibrium isotherms for 1,3-C4H6 and 1-C4H8 Agy to see the partial pressure in this isotherm ranged from 1 x 10-5 atm with sole at the scale corresponds to a concentration equal to 10 PPM. The equilibrium amounts of

The last of last amount wis somewhat small restricted absorption is the amount of 1,3-C<sub>4</sub>H<sub>6</sub> adsorbed at 150 common temperature or last amount wis somewhat small restricted absorption is the amount of the last amount of

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The lacks of 1/3-C, H, adsorbed in the low pressure reduced as regime with the decrease in temperature could be attributed as your ASAP 2010 to cut off adsorption before uptake completion as a particular pressure point if the system is too slow as a few diffusing of This possible error was corrected and a new as a certain distribution isotherm afor a 1/3-C, H, one AgY absorbte was included as per an example which follows below.

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Confirmation of Selectivity: Table 14

of the sorbent, it was tested with C2H4 at 70°C. The equilibrium isotherms of C2H4 on NaY and Agy type zeolites at 70°C are shown in Figure 5. In this figure, the effect of m-complexation can be clearly observed at low pressures. However, at higher pressures, this effect is somewhat the diminished. This phenomenon could be attributed to a filling of the pare cavity of Y-type zeolite. However, for this particular application, the behavior of the sorbent at high loadings is not important at It is the low pressure to an application.

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behavior of the sorbent which is critical for this particular purification-type application.

# Example 4

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A-Type Zeolite Selectivity for 1,3-Butadiene
The structural unit of Type A zeolite is the sodalite cage. The unit cell of Type A zeolite is the AlO<sub>4</sub> and 12 SiO<sub>4</sub> (see Figure 1). There are twelve negative charges to be balance by cations in each cell unit. The free diameter in the central cavity is 11.4 Å (Yang, 1987). Access to the pores is restricted by 8-membered oxygen rings with a free aperture of 4.3 Å in the unobstructed Ca2+ grant AE bri (5A) form. This can be reduced to 3.8 A by exchanging with Nat and to 3.0 A by exchanging with Kt. It, is this wide a vilues f variation of pore sizes and smaller cavity size that makes

type A zeolites desirable for butadiene separation.

Type A zeolites desirable for butadiene separation.

Commercially available (5A) Zeolite from Linde

Was tested: Low pressure equilibrium isotherms for 1,3-C,4H,

and 1-C,4H, on 5A at 25°C are shown in Figure 6. Equilibrium amounts of 1,3-C<sub>4</sub>H, and 1-C<sub>4</sub>H, at 1 atm were measured at 3.8 and 3:3 mmol/g, respectively. The low pressure adsorption of this material was excellent. At a concentration of 50 PPM, this material adsorbed 0.8 mmol/g of 1.3 C<sub>4</sub>H<sub>6</sub>. The working capacity of this sorbent between 50 and 10 PPM is also excellent. This was measured at approximately 0.7 25 mmol/g.

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Example 5
5A Zeolite and 5A Ion-Exchanged Zeolite Selectivity

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This example builds on the success of 5A type zeolite in adsorbing 1,3-C,H, at very low loadings. Here, 30 Athe focus was to reduce 1-C.H. adsorption. As mentioned someone maintaining is an about the some a sorbent for Restruction musically replicable 30.0 world hard of more than the control of the strategies to synthesize a sorbent for the strategies and the strategies and the strategies and the strategies and the strategies are sold to sold the strategies as the strategies and the strategies and the strategies are sold to sold the strategies and the strategies are sold to sold the strategies and the strategies are sold to sold the strategies and the strategies are sold the strategies and the strategies are sold to sold the strategies and the strategies are sold to sold the strategies and the strategies are sold to sold the strategies and the strategies are sold to sold the strategies are sold to sold the strategies and the strategies are sold to sold the strategies and the strategies are sold to sold the strategies are sold to sold the strategies and the strategies are sold to sold the strategies are sold the strategies and the strategies are sold to sold the strategies this application was to use kinetic or steric effects to separate 1,3-C4H6 (butadiene). from 1-C4H8 (butene).

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Therefore, it is possible to limit 1-C4H8 adsorption by hindering its uptake or completely excluding it from the pore structure. This was accomplished by exchanging Ca+ cations in the pores of Type A zeolite with several cations of various sizes. The pure component equilibrium isotherm for  $1,3-C_4H_6$  and  $1-C_4H_8$  on Sr exchanged 5A at 25°C are shown in Figure 7. The pressure in this isotherm ranged from 1 x 10-5 to 1 atm. Equilibrium adsorption amount for 1,3-C4H6 on Sr exchanged 5A at 25°C and 1 atm was measured at 3.1 mmol/g. Although this sorbent showed excellent capacity atmmol/g. Atthough this solvent should be all the solvent should be solvent for the solvent should be solvent solven for 1,3-C4H6 and 1-C4H8 on Sr exchanged 5A and 5A were measured at 70°C and are shown in Figure 7. The values of D/R2 for 1,3-C,H6 and 1-C,H8 on CaA were calculated at 1.07  $\times$  10<sup>2</sup> and 7.30  $\times$  10<sup>-3</sup> 1/s, respectively. Diffusion time constants were also measured on Sr exchanged 5A and were calculated at  $8.55 \times 10^{-3}$  and  $4.06 \times 10^{-3}$  1/s, respectively. Due to the larger radius of  $Sr^{2+}$  over  $Ca^{2+}$  (1.12 vs. 0.99 Å), a small reduction in 1-C4He uptake rates is observed. However, the reduction in 1-C,H, uptake is not significant enough for a kinetic separation scheme. If the ion exchange is performed with the next larger cation, Ba2+, with a radius of 1.34 A, the uptakes of both adsorbents are severely hindered. Therefore, it is questionable whether hindrance of 1-C4H, access to the pore structure of A type zeolites using simple ion exchange techniques is possible without also blocking 1,3-C4H6. The small difference in kinetic diameter between the two molecules is problematic. 30 Desorption experiments were carried out on 1,3-C4H6 on 5A zeolite at 70°C. However, these experiments were performed in the Shimadzu TG-50 TGA. Therefore, they do not include data below 0.06 atm. Equilibrium amounts of 1,3-C4H6 on 5A Zeolite at 70°C and 1 atm were measured at 2.81 mmol/g. After desorption, at 1 atm, the sample was exposed to a helium stream at 70°C to regenerate it. However, this procedure only removed 1.23 mmol/g which left

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1.58 mmol/g of  $1.3-C_4H_6$ . This final amount was removed by heating the sample to 180°C in the presence of helium. A desorption curve is shown in Figure 8. As per the above examples, success has been achieved with 5A zeolite for 1,3-butadiene separation. It is not clear yet the reason for the success of 5A and the lesser performance of Sr exchanged 5A which is very similar structurally. The pore size of this zeolite determines dits effectiveness for C6 Limit 1.7 % and C8 separation surgean sasw an radical of the

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1,3 Butadiene/1-Butene Separation: π-Complexation Effects The state of the previous examples there is shown 1,3-butadiene (C.H. adsorption on AgY zeolite at various temperatures. These examples did not include data to directly compare the  $15_{\odot}$  effects of  $\pi$ -complexation on  $C_4H_6$  adsorption. This example av. aca direct comparisonaby measuring CH, adsorption on the sodium-exchanged type Y zeolite (NaY) of It is known that the sodium cation is not capable of mecomplexating with olefins. A comparison involving AgA and CaA (5A) was not used because of the dominance of kinetic effects in the AgA system. The pore size of AgA lies in the range between 3A and 4A type zeolites.

> Equilibrium adsorption isotherms for  $C_4H_6$  on AgY and NaY at 120°C are shown in Figure 9. The effect on  $\pi$ complexation is observed in the equilibrium amounts of  $C_4H_6$ adsorbed on AgY and NaY at 1 atm which were measured at 3.3 and 2.6 mmol/g, respectively. However, the adsorption of  $C_4H_6$  at 50 PPM of saturation pressure of 1-Butene ( $C_4H_8$ ) at 120°C (1.1 x 101<sup>-3</sup> atm) is a more significant parameter. Using this parameter, the effects of  $\pi$ -complexation are The adsorption amounts of  $C_4H_6$  at 50 PPM more significant. on AgY and NaY were measured at 2.7 and 0.3 mmol/g, respectively. The working capacity of the sorbents in the range of 50 to 10 PPM (1.1 x  $10^{-3}$  -  $2.2 \times 10^{-4}$  atm) was calculated at 2.35 and 0.2 mmol/g, respectively. When the working capacities of the sorbents are calculated,

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superior performance of  $\pi$ -complexation sorbents becomes

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alog ad 5 plantaber Insthe markier examples, The isotherms of 1-

Added isotherms were measured at 70°C. The isotherms are shown in Figure 10. The amounts adsorbed are lower at 70°C. More significantly, the heel of the isotherm for butadienes is shifted towards a higher partial pressure. Fortunately, however, the amount adsorbed is quite high at a pressure butadiene partial pressure of 50°ppm of 10°atm (i.e., the add that a saturation pressure of butene at 70°C. The amount is approximately 1 mmob/go The amount adsorbed at 10 ppm of 10 atm (i.e., atm tis nearly zeros There 54 zeolite is an attractive

manned to a sorbent for this selective removal of diene at 70°C (at a guidene located pressure of 10°atm) of The polar nature of the surface env (AR) associating zepliteris though to contribute to the separation.

The analysis of an approximate analysis of the effect of the state of the end of the end

Table I.
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LSX, Y and A zeolite nominal chemical formulas

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The invention provides substantial advantages over conventional methods for separating diene from a mixture. The adsorbents and methods of the invention are useful for purifying mono-olefin from a mixture which includes mono-olefin and diene. The adsorbents and methods can be used to purify either the dienesor to purify the mono-olefin. The adsorbents and methods as exemplified in the examples demonstrate particular effectiveness removing a diene from a mixture comprising the diene and one or more mono-olefins. The examples show effective separation results using gas-phase separation at the pressures and temperatures stated therein. The adsorbents and processes are demonstrated for a saturated 1-butene (olefin) stream, which is contaminated with 1,3-butadiene (diene), typically at one weight percent or less. separation in this case may include both gas and liquid phase selective adsorption due to the saturated nature of the 1-butene olefin stream. The processes described herein are described with reference to separation of C4 to C8 diene from a C4 to C8 mono-olefin. The physical characteristics of the adsorbents result in very effective separation as demonstrated by the examples. the chosen conditions, the separation is accomplished in the gas phase, the liquid phase or some combination thereof, including, for example, saturated gas. reasons described above, the adsorbents are also considered to be useful for separation of other dienes, that is C4 to C10 dienes from C4 to C10 mono-olefins. Beyond C10 the process is also thought to be applicable, for example, C12 diene/C12 mono-olefin separation. However, the selectivity and economic benefits are expected to decline for the higher hydrocarbon mono-olefin/diene separation.

> The invention provides good results at operating temperatures and pressures that are not extreme. results are achievable at pressures on the order of subatmospheric (i.e., 0.01 to 0.1 atm) to moderately high (i.e. 20 to 30 atm) and in a pressure swing adsorption

method. Good results are achievable at temperatures from ambient or a desired feed temperature (i.e. 0 to 25°C) and up to moderately high (i.e. 250°C), and in a temperature swing adsorption method. With the adsorbents of the invention, temperature swing or pressure swing adsorption processes are usable for effective separation between dienes and mono-olefins. This is demonstrated by the very effective purification of butenes, by removing 1,3-butadiene at concentrations above 20 ppm to final concentration below 10 ppm. Therefore, the invention provides a very attractive solution to the problem of selective adsorption of trace amounts of diene to separate diene from a mixture

of certain embodiments thereof, it is not intended that it be limited to the above description, but rather only to the extent set forth in the following claims.

The embodiments of the invention in which an exclusive property or privileged is claimed, are defined in the following claims.

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1. A process for separating a diene from a mixture including said diene; said process comprising the  $\mathbb{F}[q]$  . The section  $\mathbf{f}$  is the  $\mathbf{f}$  and  $\mathbf{f}$  is a section of  $\mathbf{f}$  and  $\mathbf{f}$  is  $\mathbf{f}$  and  $\mathbf{f}$  and  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  are  $\mathbf{f}$  and  $\mathbf{f}$  are  $\mathbf{f}$ contacting said mixture with an adsorbent which preferentially adsorbs said diene, thereby producing a nonadsorbed component and aldene rich adsorbed component, wherein said adsorbent comprises at least one of (a) and 1 3 (b) : tydiega, at congenimatique spova, 20noise avei sale , each (a) an ion-exchanged zeolite selected from the ic melders end to mois group consisting of zeolite X, zeolite Y, odrigate or could continue and zeolite LSX, said zeolite exchangeable cationic sites, and emmed willedimmed and maintains of said sites having JI LENT FORESTIL CONTROL CATION OF COPPER CATION PRESENT; ed a contract the contract of the faving exchangeable cationic sites, a plurality of said A zeolite sites as spine of orlineval educating at an alkalite metal cation

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2. The process of claim 1 wherein said mixture comprises at least one mono-olefin having as many carbon atoms as said diene.

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- 3. The process of claim 2 wherein said diene is selected from the group consisting of butadiene, hexadiene, octadiene and mixtures thereof; and said mono-olefin is selected from the group consisting of butene, hexene, octene and mixtures thereof.
- 4. The process of claim 2 wherein said monoolefin is butene and said diene is butadiene.
- 5. The process of claim 2 wherein said monoolefin is hexene and said diene is hexadiene.

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- 6. The process of claim 2 wherein said monoolefin is octene and said diene is octadiene.
- 7. The process of claim 2 wherein said mixture comprises said mono-olefin in a gaseous state and saturated with said diene.
- 8. The process of claim 1 wherein said type LSX zeolite has a silicon to aluminum (Si/Al) atomic ratio of less than or equal to 1.2.
- True s 5. s (3) maining ai dollars facts illes 9. The process of claim 8 wherein said ratio is one.
- 10. The process of claim 1 wherein said X and Y zeolites respectively have a silicon to aluminum ratio of 1:1 and 1000:1.
- 11. The process of claim 1 wherein at least half of the cationic sites of said ion-exchanged zeolite contain said silver cation.
- 12. The process of claim 1 wherein essentially all cationic sites of said ion-exchanged zeolite contain said silver cation.
- 13. The process of claim 1 wherein at least half of the cationic sites of said ion-exchanged zeolite contain said copper cation.
- 14. The process of claim 1 wherein essentially all cationic sites of said ion-exchanged zeolite contain said copper cation.
- The process of claim 1 wherein said type A zeolite has the nominal general formula  $M^{+1}_{x}Z^{+2}A^{+3}_{b}[(AlO_{2})_{12}]$  (SiO<sub>2</sub>)<sub>12</sub>] wherein  $M^{+1}$  is an alkali metal cation,  $Z^{+2}$  is an

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alkaline earth metal cation, and  $A^{+3}$  is a tri-valent cation,  $0 \le x \le 12$ ,  $0 \le y \le 6$ , and  $0 \le b < 4$  provided that: x + 2y + 3b is 12.

- 16. The process of claim 15 wherein said A<sup>+3</sup> is a metal cation and said zeolite comprises said A<sup>+3</sup> in an a atomic amount less than the combined atomic amount of said alkali metal cation and said alkaline earth metal cation.
  - 17. The process of claim 15 wherein the alkaline earth metal cation is calcium (Ca) and at least half of the cationic sites contain calcium.
  - 18. A process of separating a diene selected from the group consisting of butadiene, hexadiene, octadiene and mixtures thereof from a mixture including said diene, said process comprising the steps of:

contacting said mixture with an adsorbent which preferentially adsorbs and releasably retains said diene at a selected temperature and pressure, thereby producing a non-adsorbed component and a diene-rich adsorbed component, wherein said adsorbent comprises at least one of (a) and (b)

- (a) an ion-exchanged zeolite selected from the group consisting of zeolite X, zeolite Y, and zeolite LSX, said zeolite having cation exchange sites, a plurality of said sites having silver cation or copper cation present;
- (b) type A zeolite having cation exchange sites, a plurality of said sites having an alkali metal cation or an alkaline earth metal cation present;

and then changing at least one of said pressure and temperature to thereby release said diene-rich component from said adsorbent.

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pressure of preferential adsorption is a first pressure, and the pressure of release is a second pressure less than said first pressure.

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The process of claim 19 wherein said second process of about 0.01 atmosphere to about 5 atmospheres.

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The process of claim 19 wherein said second pressure is in a range of about 0.1 atmosphere to about 0.5 atmospheres.

33. The process of claim of wherean samathelity

temperature of preferential adsorption is a first temperature, and the temperature of release is a second temperature greater than said first temperature.

The recommendation of the processof of a community wherein said first standard the first standard the process of about 0°C to about 150°C.

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temperature is in a range of about 25°C to about 80°C.

Landron end a sister 27:27 The process of claim 24 wherein said second of ACLAD we temperature is in a range of about 70°C to about 250°C.
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moisso smells 128. The process of claim 24 wherein said second of the process of about 100°C to about 120°C.

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A process for separating 1,3-butadiene from the state of comprising 1,3-butadiene and at least one other of contacting said mixture with an adsorbent which preferentially adsorbs said 1,3-butadiene, said adsorbent which comprising a silver of copper ion-exchanged zeolite is selected from the group consisting of Y zeolite, X zeolite, low silica X zeolite (LSX), and combinations thereof.

Lania libra misser \$2.30.5. The process of Claim 29 wherein said zeolite and the comprises a plurality of Cation exchangeable sites and at least half of said sites contain silver cations.

Boot a pass of wall of said cationic sites contain silver cations.

32. The process of claim 29 wherein said zeolite bronse bise comprises amplification exchangeable sites and at 2.0 zoods pleast half of said sites confain copper cations.

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33. The process of claim 32 wherein essentially bedoels establiofisate cathonic sites contain said copper cations.

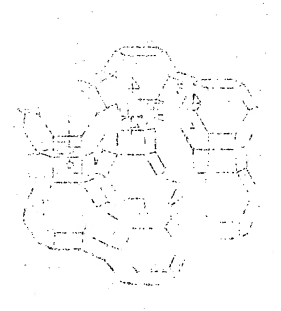
The process for separating 1,3 but addene and at Teast one other C4 unsaturated compound, said process comprising the step of the officing said mixture with an adsorbent which the preferentially adsorbe said 1,3 but addene, said adsorbent comprising a type A zeolite having cation exchange sites, said as a salurality of said sites having an alkali metal cation or the same alkaline earth metal cation.

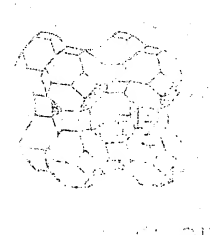
booker of the last of 35% to The opposess of claim 34 wherein the nominal content of sugmeral formulas of said zeolite list  $M^{+1}_{x}Z^{+2}_{y}A^{+3}_{b}[(AlO_{2})_{12}]$  (SiO<sub>2</sub>)<sub>12</sub>] wherein  $M^{+1}$  is an alkali metal cation,  $Z^{+2}$  is an time alkaline earth cation, and  $A^{f3}$  is a tri-valent cation, and  $A^{f3}$  is a tri-valent cation,

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6, and the value of b is at minimum zero and at maximum less than the sum of x plus y, provided that: x + 2y + 3b is 12.

36. The process of claim 34 wherein the alkaline earth cation is calcium (Ca) and more than half of the cation exchange sites contain calcium.





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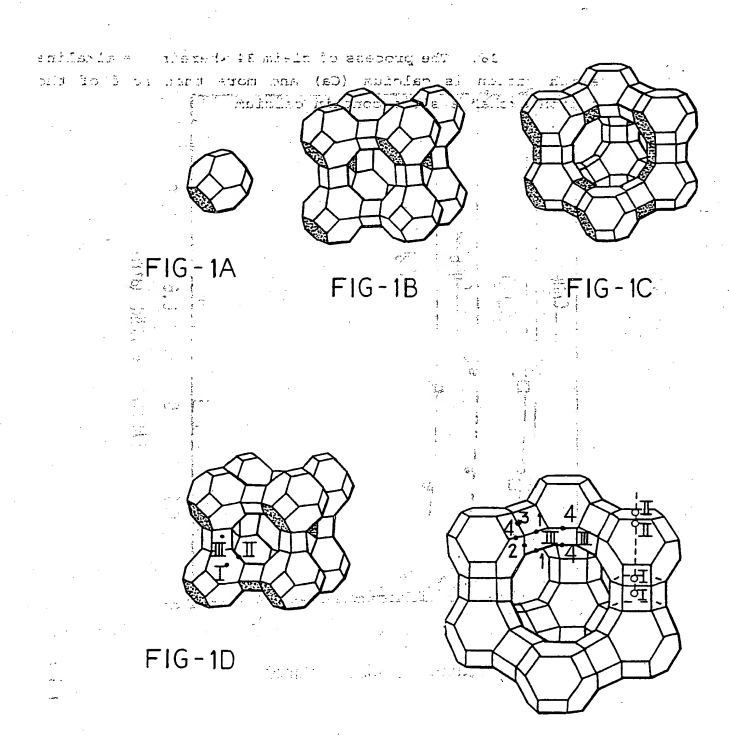
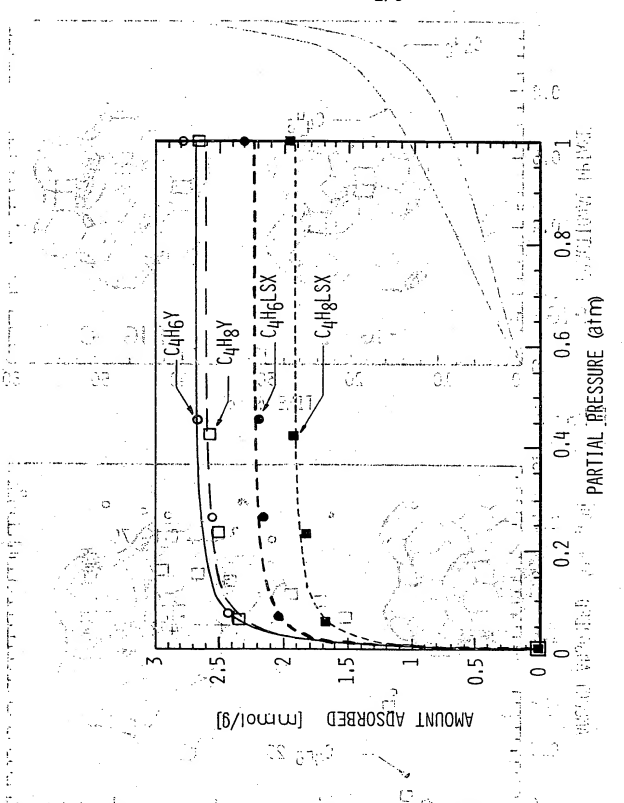


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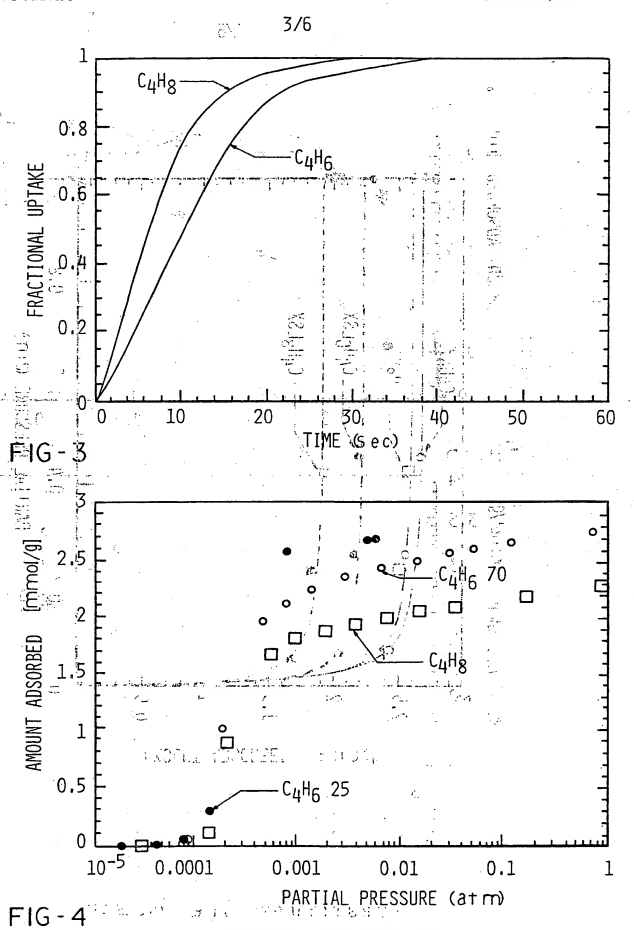
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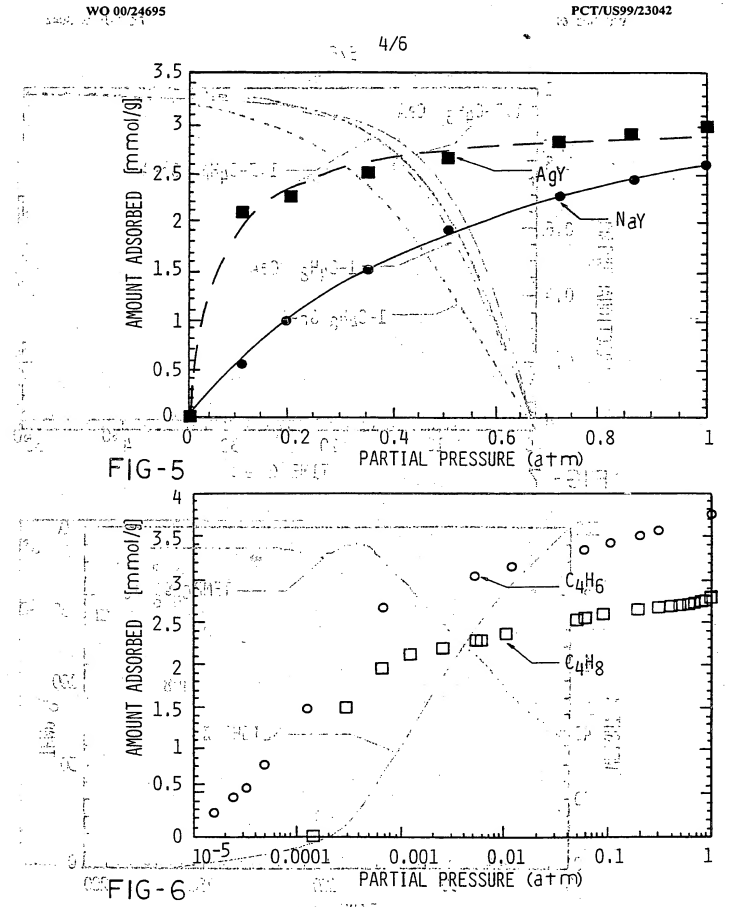
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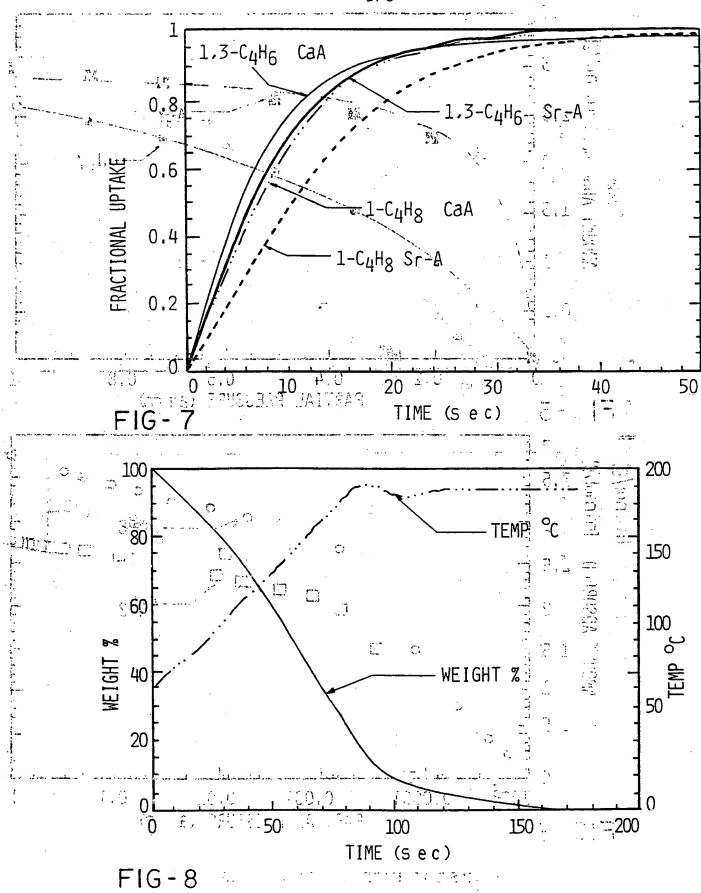
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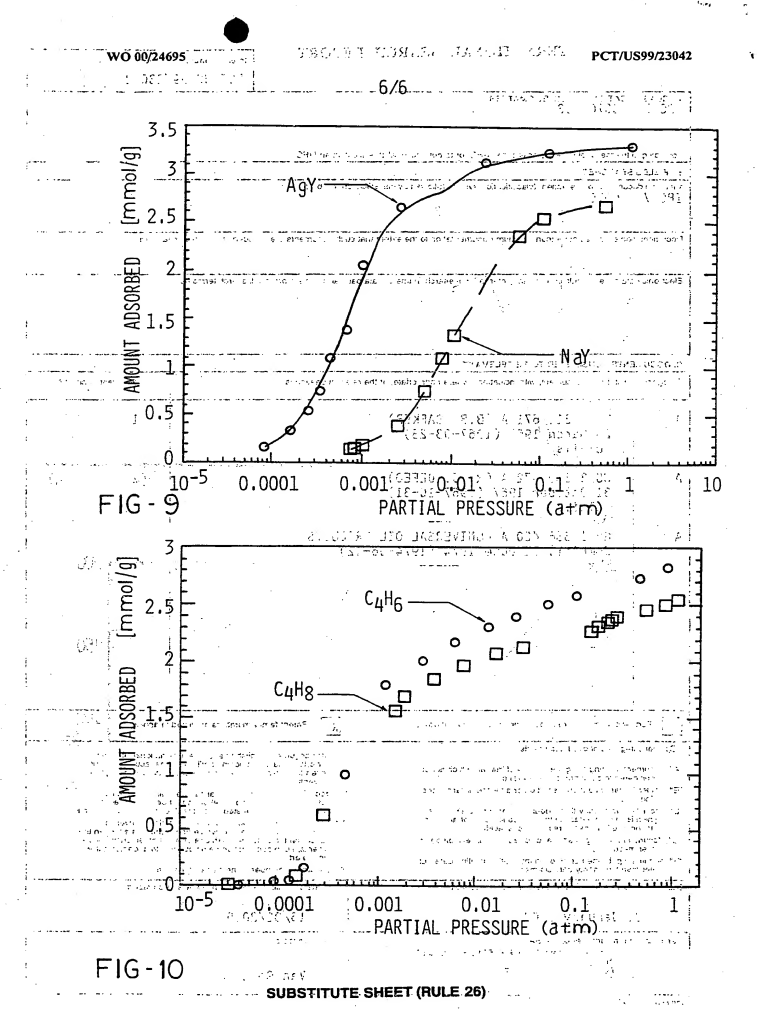


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# publication in the search report

Inter anal Application No

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	C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	e.		<u> </u>			
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